Chemistry of Epoxy Compounds. 'V.2 Preparation of Some Hydroxy-ethers from 9,10-Epoxystearic Acid and 9,10-Epoxyoctadecanol

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The reaction of oxirane compounds with compounds containing a hydroxyl group offers a convenient way of preparing many types of hydroxyethers (Equation 1).

Acidic or alkaline materials are usually employed to catalyze this reaction, although for the more reactive oxirane compounds, a catalyst is not necessary. The reaction is generally applicable, and numerous hydroxy-ethers which contain a variety of functional groups can be readily prepared. Most of the published work has described the reaction of short-chain oxirane compounds with saturated aliphatic alcohols, and little information is available on the reaction of long-chain oxirane compounds with hydroxyl compounds. In this paper, the preparation of some hydroxy-ethers by the reaction of 9,10-epoxystearic acid and 9,10epoxyoctadecanol with methyl, ethyl, n-propyl, n-butyl, iso-butyl, n-octadecyl, allyl, and β -chloroallyl alcohol and phenol, respectively, is reported. (Mixtures of products are obtained, since it is equally probable that the oxirane ring is opened between the oxygen atom and C₀ or C₁₀.)

The reaction of 9,10-epoxystearic acid (prepared from oleic acid by hypochlorination followed by dehydrohalogenation) with a large excess of ethyl, n-propyl and n-butyl alcohol, respectively, in the presence of catalytic quantities of sulfuric acid, has been reported by Nicolet and Poulter.3 The reaction products which they obtained were saponified with aqueous alkali, and the three compounds isolated were mixtures of the 9- and 10monoalkyl ethers of 9,10-dihydroxystearic acid. Yields were not reported. The oxirane ring of 9,10-epoxystearic acid can best be opened by heating the acid with a large molar excess of the alcohol and catalytic quantities of sulfuric acid, and under these conditions the carboxyl group is quantitatively esterified. By omitting the saponification step we obtained the hydroxy-ethers shown in Table I, in which the alkyl radicals in the ester and ether groups are identical. These products were isolated by vacuum distillation of the re-

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(2) For previous papers, see THIS JOURNAL, 66, 1925 (1944);

67, 412, 1786 (1945); 68, 1504 (1946).

action products. We also treated 9,10-epoxystearic acid with n-octadecyl alcohol, phenol and β-chloroallyl alcohol under the conditions described above in an attempt to prepare n-octadecyl 9,10(10,9)-octadecoxyhydroxystearate, 9,10(10,-9)-phenoxyhydroxystearic acid, and β -chloroally-9,10(10,9)-chloroalloxyhydroxystearate, respectively, but we were unable to purify the products completely. The analytical data which we obtained indicated, however, that in the main the reaction had proceeded in the expected manner.

We were unable to find any literature references to the reaction of 9,10-epoxyoctadecanol with hydroxyl compounds. We prepared the compounds shown in Table II in 25 to 65% yields, by heating 9,10-epoxyoctadecanol with a large molar excess of the appropriate alcohol in the presence of sulfuric acid as catalyst, and subsequently distilling the reaction products. We were unable to isolate pure 9,10(10,9)-n-octadecoxyhydroxyoctadecanol, 9,10(10,9)-phenoxyhydroxyoctadecanol and 9,10(10,9)-β-chloroalloxyhydroxyoctadecanol, although as mentioned earlier, the reactions proceeded in the expected manner.

The purified hydroxy-ethers are colorless, odorless, high-boiling liquids, soluble in all the common organic solvents and insoluble in water. They do not solidify when stored at 0° for long periods of time. Yields and characteristics of the products are shown in Tables I and II. Their high boiling points, insolubility in water, and stability suggest their possible use as plasticizers.

Allyl 9,10(10,9)-alloxyhydroxystearate (Table I) was co-polymerized in various proportions (from 1 to 40% by weight of total monomers) with vinyl acetate, according to the procedure of Guile and Huston.4 Over the entire percentage range, insoluble, infusible products were obtained which ranged in physical appearance from hard, glasslike resins to soft, rubbery gels, thereby indicating that cross-linking probably had occurred and that both allyl groups must have reacted. That the fatty derivative was probably completely chemically bound was demonstrated by the fact that no oil was extruded when the co-polymers were squeezed in thin layers between sheets of absorbent paper.

In the reactions of 9,10-epoxystearic acid with hydroxyl compounds, small quantities of byproducts, tentatively identified as esters of 9,10dihydroxystearic acid,5 were formed, and in the re-

⁽³⁾ Nicolet and Poulter, ibid., 52, 1186 (1930). These workers gave 53.8° as the melting point of the 9,10-epoxystearic acid which they employed. It has been shown by Ellis, Biochem. J., 30, 753 (1936), and confirmed in our laboratory, that the correct melting point of 9,10-epoxystearic acid prepared from oleic acid by hypochlorination followed by dehydrohalogenation is 59.5°.

⁽⁴⁾ Guile and Huston, "A Revised Laboratory Manual of Synthetic Plastics and Resinous Materials," Michigan State College, 1944, p. 99.

⁽⁵⁾ Swern and Jordan, This Journal, 67, 902 (1945); Swern Jordan and Knight, ibid., 68, 1673 (1946).

R	Formula	Yield,	Boiling range °C. Mm.		fication equivalent Calcd, Found		Hydroxyl, % Calcd. Found		Carbon, %b Calcd. Found		Hydrogen, %b iCalcd. Found		n ⁸⁰ D	Molec refrac d^{∞}_4 Calcd.		ction	
Methyl	C20H40O4	85	186-189	0.7-1.1	344.5	344.7	4.94	4.95	69.7	69.9	11.7	11.8	1.4500	0.9306	99.4	99.5	
Ethvl	C22H44O4	50	160-190	.02-0.2	372.6	368.2	4.56	4.57	71.0	70.7	11.9	12.0	1.4479	.9154	108.6	108.9	
n-Propyl	C24H48O4	45	180-184	.02-0.04	400.7	399.5	4.25	4.20	72.0	72.1	12.1	12.0	1.4490	.9099	117.9	118.1	
n-Butyl	C26H52O4	45	185-190	.01	428.7	425.2	3.97	3.95	72.9	72.6	12.2	12.2	1.4498	.9051	127.1	127.1	
iso-Butyl	C26H62O4	40	172-180	.02-0.04	428.7	427.2		3.76					1.4471	.8989	127.1	127.5	
Allyle	C24H44O4	60	196-202	.35-0.5	396.6	387.5	4.59	4.71	72.7	72.2	11.2	11.2	1.4589	.9266	116.9	117.1	

^a Purified products, after at least two distillations. ^b Analyses by Mary J. Welsh of this Laboratory. ^c Iodine number: calcd., 128.0; found, 126.0.

Yield. Boiling range					xvl. %	Carbon, %b		Hydrogen, %b				Molecular refraction			
Formula	%ª	°C.	Mm.	Calcd.	Found	Calcd.	Found	Calcd.	Found	n ³⁰ D	d ²⁰ 4	Calcd.	Found		
C19H40Os	65	158-161	0.007-0.008	10.7	10.8	72.1	71.5	12.7	12.6	1.4584	0.9151	94.6	94.5		
C20H42O2	60	179-185	.005-0.01	10.3	9.82	72.7	73.0	12.8	12.5	1.4565	.9078	99.3	99.7		
CnH4O	30	156-168	.005-0.01	9.87	9.32	73.2	73.1	12.9	12.7	1.4559	.9040	103.9	103.6		
C22H46O1	50	170-175	.007	9.49	9.42	73.7	73.5	12.9	13.1	1.4562	.9002	108.5	108.3		
C22H46O2	25	163-170	.007-0.02	9.49	9.29	73.7	73.8	12.9	13.0	1.4550	.8980	108.5	108.3		
C21H42O2	40	171-188	.02-0.03	9.93	9.83	73.6	73.4	12.4	12.3	1.4627	.9166	103.4	102.9		
	C19H40O2 C20H42O2 C21H4O2 C22H46O2 C22H46O3	Formula % ^a C19H40Ot 65 C20H42Ot 60 C21H44Ot 30 C22H45Ot 50 C22H46Ot 25	Formula % °C. C19H40O1 65 158-161 C20H41O2 60 179-185 C21H44O2 30 156-168 C11H45O1 50 170-175 C22H46O2 25 163-170	Formula % ⁴ °C. Mm. C19H40O1 65 158-161 0.007-0.008 C20H43O2 60 179-185 .005-0.01 C11H44O1 30 156-188 .005-0.01 C11H45O1 50 170-175 .007 C12H44O1 25 163-170 .007-0.02	C19H40O: 65 158-161 0.007-0.008 10.7 C20H4:O: 60 179-185 .005-0.01 10.3 C31H4:O: 30 156-168 .005-0.01 9.87 C31H4:O: 50 170-175 .007 9.49 C31H4:O: 25 163-170 .007-0.02 9.49	C19H40Ot 65 158-161 0.007-0.008 10.7 10.8 C20H44Ot 60 179-185 .005-0.01 10.3 9.82 C31H44Ot 30 156-168 .005-0.01 9.87 9.32 C31H46Ot 50 170-175 .007 9.49 9.42 C32H46Ot 25 163-170 .007-0.02 9.49 9.29	C19H40O: 65 158-161 0.007-0.008 10.7 10.8 72.1 C20H4:O: 60 179-185 .005-0.01 10.3 9.82 72.7 C3:H4:O: 30 156-168 .005-0.01 9.87 9.32 73.2 C3:H4:O: 50 170-175 .007 9.49 9.42 73.7 C3:H4:O: 25 163-170 .007-0.02 9.49 9.29 73.7	C19H40O: 65 158-161 0.007-0.008 10.7 10.8 72.1 71.5 C20H4:O: 60 179-185 .005-0.01 10.3 9.82 72.7 73.0 C3:H4:O: 30 156-168 .005-0.01 9.87 9.32 73.2 73.1 C3:H4:O: 50 170-175 .007 9.49 9.42 73.7 73.5 C3:H4:O: 25 163-170 .007-0.02 9.49 9.29 73.7 73.8	C19H40O: 65 158-161 0.007-0.008 10.7 10.8 72.1 71.5 12.7 C20H4:O: 60 179-185 .005-0.01 10.3 9.82 72.7 73.0 12.8 C3:H4:O: 30 156-168 .005-0.01 9.87 9.32 73.2 73.1 12.9 C3:H4:O: 50 170-175 .007 9.49 9.42 73.7 73.5 12.9 C3:H4:O: 25 163-170 .007-0.02 9.49 9.29 73.7 73.8 12.9	Formula	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Formula Yield So Boiling range Hydroxyl, % Carbon, %b Calcd. Found Found Calcd. Found Calcd. Found Found Calcd. Found Found Calcd. Found Found Calcd. Found Calcd. Found Found Calcd. Found Found Calcd. Found Found Calcd. Found Found Found Calcd. Found Found Found Found Found Calcd. Found Fou	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	

^e Purified products, after at least two distillations. ^b Analyses by Mary J. Welsh of this Laboratory. ^e Iodine number: calcd. 74.1; found, 73.3.

actions of 9,10-epoxyoctadecanol, the by-product was 9,10-dihydroxyoctadecanol. These side-reaction products were separated from the hydroxyethers by crystallization from acetone.

Experimental

Starting Materials.—9,10-Epoxystearic acid, m. p. 59.5°, and 9,10-epoxyoctadecanol, m. p. 54°, were prepared by epoxidation of pure oleic acid and oleyl alcohol, respectively.² The short-chain aliphatic alcohols were the purest commercial grades, and they were distilled through an efficient fractionating column (40 to 50 theoretical plates) before use. The n-octadecyl alcohol, m. p. 57-58°, was the Eastman Kodak purest grade. The phenol was the U. S. P. grade.

Preparation of Hydroxy-ethers from 9,10-Epoxystearic Acid.—A typical reaction procedure suitable for the shortchain alcohols is given: 9,10-Epoxystearic acid (29.9 g., 0.1 mole) was dissolved in the freshly distilled anhydrous alcohol (4 ml. per gram), with gentle heating. The solution was cooled below 30°, and 0.3 g. of 95% sulfuric acid was added dropwise while the reaction mixture was shaken by hand. A temperature rise of several degrees was always observed during the addition of the catalyst. The always observed during the addition of the catalyst. solution was heated on the steam-bath for two hours, and a quantity of sodium bicarbonate equivalent to the sulfuric acid was then added. The reaction mixture was fractionally distilled, first at atmospheric pressure to recover unreacted alcohol and then under high vacuum to isolate the hydroxy-ether. A short Vigreux column was employed during the vacuum distillation, which was carried out as rapidly as possible, otherwise some debydration of the product occurred. The hydroxy-ether was dissolved in acetone (2 to 3 ml. per gram), and the solution was cooled to -20° to precipitate the ester of 9,10-dihydroxystearic acid, which was always obtained as a by-product. This was separated by filtration and rejected. The acetone was evaporated from the filtrate, and the residual oil consisted of almost pure hydroxy-ether without further treatment (yield 60 to 95%). This residue was rapidly redistilled, and fractions having the same refractive index were combined for analysis. The results are summarized in Table I.

Preparation of Hydroxy-Ethers from 9,10-Epoxyocta-decanol.—These were prepared and isolated as described under the preparation of hydroxy-ethers from 9,10-epoxystearic acid. The results are summarized in Table II.

Benzoyl Peroxide-Catalyzed Co-polymerization of Allyl 9,10(10,9)-Alloxyhydroxystearate with Vinyl Acetate.— This was conducted as recommended by Guile and Huston, a maximum content of allyl 9,10(10,9)-alloxyhydroxystearate of 40% by weight of the total monomers and 0.5% by weight of benzoyl peroxide being used. Solubility tests on the co-polymers were conducted in acetone at the boiling point and in acetic acid and amyl acetate at 100°. When thin layers of the co-polymers were squeezed between sheets of absorbent paper, no extrusion of oil was observed.

Summary

Twelve hydroxy-ethers have been prepared from 9,10-epoxystearic acid and 9,10-epoxyoctadecanol by reaction with methyl, ethyl, n-propyl, n-butyl, isobutyl and allyl alcohol, respectively. The high boiling points of the products, their insolubility in water and their stability suggest their possible use as plasticizers.

The benzoyl peroxide-catalyzed co-polymerization of allyl 9,10(10,9)-alloxyhydroxystearate (prepared from 9,10-epoxystearic acid and allyl alcohol) with vinyl acetate has been studied. Over the entire percentage range investigated (allyl 9,10(10,9)-alloxyhydroxystearate content from 1 to 40% by weight of total monomers), insoluble and infusible co-polymers which ranged from hard, glass-like resins to rubbery gels were obtained, thereby indicating that cross-linking probably occurred and that both allyl groups take part in the co-polymerization reaction.

The reaction of 9,10-epoxystearic acid and 9,10-epoxyoctadecanol with n-octadecyl alcohol,

phenol and β -chloroallyl alcohol was also investigated. Although pure hydroxy-ethers were not obtained, analytical data indicated that

the reaction proceeded mainly in the expected manner.
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